chromosome. This locus is depicted in Figures 2A and 3. Since everninomic in is only known to be produced in M. carbonacea, for the sake of particularity the EV biosynthetic pathway is associated with this microorganism. However, it should be understood that this term encompasses EV biosynthetic enzymes (and genes encoding such enzymes) isolated from any M. carbonacea, and furthermore that these genes may have novel homologues in related actinomycete bacteria that fall within the scope of the claims here. In specific embodiments, these genes are depicted in Figure 11 (SEQ ID NO:1; open reading frames and polypeptides designated as SEQ ID NOS: 2-175) and Figure 12 (SEQ ID NO: 182; open reading frames and polypeptides designated as SEQ ID NOS: 183-204). It is noted that the sequences of Figures 11 and 12 are linked (contiguous) or connected such that they are part of the same cluster, i.e., the sequence in Figure 12 precedes that of Figure 11. Moreover, the present inventors have identified specific categories into which many of the genes from the EV biosynthetic pathway fall, including but by no means limited to, orsellinic acid biosynthetic enzymes, sugar biosynthetic enzymes, glycosyltransferases, tailoring enzymes, regulatory enzymes (serinethreonine kinases), and resistance mechanism enzymes (rRNA methylases and transporter enzymes). These categories are discussed in greater detail, infra. The gene products are listed in Tables 1a and 1b.

Table 1a. Gene Products and Putative Enzymatic Functions
Involved in Everninomicin Production

| Gene<br>Product         | CDS <sup>1</sup> | RBS <sup>2</sup> | SEQ ID NO. | Enzymatic Function (Protein ACC No; BLAST Score)                                | Class                 |
|-------------------------|------------------|------------------|------------|---|-----------------------|
| evdA<br>length<br>416aa | (1321382)*       | (13891394)*      | 2, 3       | similarity to hydroxylase<br>(CAA11782; 6.5e-137)                               | sugar<br>biosynthetic |
| evdB<br>length<br>373aa | (14902611)*      | (26182622)*      | 4, 5       | hexose aminotransferase,<br>dnrJ homolog<br>(daunorubicin)<br>(P25048; 2.8e-65) | sugar NH2<br>addition |
| evdC<br>length<br>412aa | (26223860)*      | (38673870)*      | 6, 7       | similar to flavoprotein,<br>oxidase<br>(S39965; 4.4e-92)                        | sugar<br>biosynthetic |
| evdD<br>length<br>389aa | (41435312)       | (41344138)       | 8, 9       | dNTP-hexose<br>glycosyltransferase<br>(AAC01731; 4.6e-49)                       | Glycosyl<br>transfer  |
| evdE<br>length<br>308aa | (53096235)       |                  | 10, 11     | hexose dehydratase<br>(CAA18814; 8.0e-58)                                       | sugar<br>biosynthetic |
| evdF<br>length<br>347aa | (62327275)       | (62266229)       | 12, 13     | dNTP-hexose<br>glycosyltransferase<br>(CAB07092; 3.4e-18)                       | Glycosyl<br>transfer  |
| evdG<br>length<br>351aa | (72728327)       |                  | 14, 15     | unknown .   | unknown               |

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|                          |                 |                 | -8-    |   |   |
|--------------------------|-----------------|-----------------|--------|---|---|
| evdH<br>length           | (83429364)      | (83338336)      | 16, 17 | dNTP-hexose<br>glycosyltransferase  | Glycosyl<br>transfer                    |
| 340aa<br>evdI            | (946310,224)*   | (10,23210,235)* | 18, 19 | (CAA19930; 0.8)<br>hydrolase  | sugar                                   |
| length<br>253aa          |                 |                 |        | (AAB81835; 6.8e-10)   | biosynthetic                            |
| evdJ<br>length<br>250aa  | (10,42411,176)  |                 | 20, 21 | unknown   | unknown                                 |
| evdK<br>length<br>415aa  | (11,20812,455)  |                 | 22, 23 | hexose dehydratase or<br>empimerase<br>(CAB08849; 3.3e-26)                            | sugar<br>biosynthetic                   |
| evdL<br>length<br>304aa  | (12,10813,022)* | (13,02713,030)* | 24, 25 | dNTP-hexose<br>glycosyltransferase<br>(S37028; 0.010)                                 | Glycosyl<br>transfer                    |
| evrA<br>length<br>317aa  | (14,41015,363)* | (15,36915,373)* | 26, 27 | hexose epimerase<br>(CAA12010.1; 1.3e-40)   | sugar<br>biosynthetic                   |
| evrB<br>length<br>344aa  | (15,38016,414)* |                 | 28, 29 | hexose oxidoreductase<br>(ACC01734; 1.3e-65)  | sugar<br>biosynthetic                   |
| evrC<br>length<br>484aa  | (16,41917,873)* |                 | 30, 31 | hexose dehydratase<br>(CAA12009; 2.2e-107)  | sugar<br>biosynthetic                   |
| evrD<br>length<br>354aa  | (17,87018,934)* |                 | 32, 33 | GDP-mannose 4,6-<br>dehydratase<br>(BAA16585; 1.0e-88)                                | sugar<br>biosynthetic                   |
| evrE<br>length<br>510aa  | (19,37420,906)  | ·               | 34, 35 | multidrug efflux<br>transporter<br>(CAB15277; 1.4e-59)                                | resistance<br>mechanism                 |
| evrF<br>length<br>492aa  | (21,06422,542)  | (21,05622,542)  | 36, 37 | similar to non-heme<br>oxygenate/halogenase<br>(CAA11780; 4.3e-58)                    | orsellinic<br>acid chlorine<br>addition |
| evrG<br>length<br>474aa  | (22,74824,172)  | (22,73622,740)  | 38, 39 | oxidase<br>(Q12737; 5.5e-67)  | tailoring                               |
| evrH<br>length<br>348aa  | (24,17725,223)* | (25,23025,233)* | 40, 41 | unknown<br>(AAB89073; 3.2e-6)   | unknown                                 |
| evrI<br>length<br>358aa  | (25,55026,626)  |                 | 42, 43 | acyl starter unit fidelity<br>(daunorubicin homology)<br>(AAA65208; 5.7e-56)          | PKS acyl<br>Carbon<br>choice            |
| evrJ<br>length<br>1264aa | (26,68530,479)  | (26,67226,676)  | 44, 45 | orsellinic acid synthase 6-<br>methylsalicilic acid<br>synthetase<br>(CAA72713; 0.0e) | polyketide<br>synthetase                |
| evrK<br>length<br>439aa  | (30,55731,876)* | (31,88531,888)* | 46, 47 | Na/H antiporter<br>(BAA16991; 2.1e-14)  | unknown                                 |
| evrL<br>length<br>313aa  | (31,94132,882)* |                 | 48, 49 | similar to gene essential to<br>heme biosynthesis<br>(BAA12681; 0.0012)               | unknown                                 |
| evrM<br>length<br>412aa  | (33,16734,405)* | (34,41434,418)* | 50, 51 | similar to p450<br>hydroxylase<br>(S18530; 3.8e-70)                                   | tailoring                               |

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|                         |                 |                 | -9-    |   |   |
|-------------------------|-----------------|-----------------|--------|---|---|
| evrN<br>length<br>253aa | (34,44935,210)* | (35,21935,221)* | 52, 53 | methyl transferase<br>(CAB10751; 0.00061)                                     | tailoring                                   |
| evrO<br>length<br>314aa | (35,29436,238)* |                 | 54, 55 | unknown<br>(BAA20094; 0.56)   | unknown                                     |
| evrP<br>length<br>242aa | (36,23536,963)* |                 | 56, 57 | unknown<br>(CAB05421; 0.00020)  | unknown                                     |
| evrQ<br>length<br>342aa | (36,99838,026)* |                 | 58, 59 | similar to oxidoreductase<br>and heat stress protein<br>(P80874; 7.8e-31)     | tailoring                                   |
| evrR<br>length<br>164aa | (38,07238,566)* |                 | 60, 61 | low similarity to hexaheme<br>nitrite reductase regulator<br>(P30866; 0.0034) | regulatory<br>(methyl<br>transferase)       |
| evrS<br>length<br>423aa | (38,89240,163)* |                 | 62, 63 | dNTP-hexose<br>glycosyltransferase<br>(AAD15267; 1.9e-36)                     | Glycosyl<br>transfer                        |
| evrT<br>length<br>224aa | (40,21640,890)* | (40,89940,902)* | 64, 65 | similar to L-proline<br>hydroxylase<br>(BAA 20094; 5.5e-7)                    | tailoring                                   |
| evrU<br>length<br>229aa | (40,88741,576)* |                 | 66, 67 | methyltransferase<br>(CAB02029; 5.6e-6)                                       | tailoring                                   |
| evrV<br>length<br>342aa | (41,67942,707)* | (42,71442,717)* | 68, 69 | dTDP-glucose epimerase<br>(AAB84886; 3.5e-36)                                 | L-dTDP-<br>glucose<br>biosynthetic          |
| evrW<br>length<br>329aa | (42,81043,799)* | (43,80743,811)* | 70, 71 | dTDP-glucose dehydratase<br>(CAA72715; 5.1e-136)                              | D-dTDP-<br>glucose<br>biosynthetic<br>(GDH) |
| evrX<br>length<br>355aa | (43,79944,866)* |                 | 72, 73 | dTDP-glucose synthetase<br>(A26984; 1.2e-118)                                 | D-dTDP-<br>glucose<br>biosynthetic          |
| evrY<br>length<br>248aa | (45,01445,760)* | (45,76745,770)* | 74, 75 | dehalogenase<br>(P24069; 5.8e-8)  | drug<br>resistance                          |
| evrZ<br>length<br>250aa | (45,96246,714)* | (45,95245,956)* | 76, 77 | similar to<br>muramidase/lysozyme<br>(P25310; 1.2e-77)                        | drug<br>resistance                          |
| evsA<br>length<br>692aa | (47,15649,234)* |                 | 78, 79 | serine threonine kinase<br>(BAA32455; 2.0e-76)                                | regulatory                                  |
| evsB<br>length<br>362aa | (51,62752,715)  | (51,62051,622)  | 80, 81 | similar to proteases  | unknown                                     |
| evsC<br>length<br>222aa | (52,88953,557)  |                 | 82, 83 | similar to MAF involved in<br>septum formation<br>(BAA18425; 1.3e-21)         | unknown                                     |
| evbA<br>length<br>217aa | (53,55454,207)  |                 | 84, 85 | O-methyl transferase<br>(AAC44130; 8.6e-38)                                   | tailoring;<br>possible<br>resistance        |
| evbB<br>length<br>251aa | (54,36255,117)* | (55,12555,128)* | 86, 87 | membrane pump, homolog<br>mithramicin resistance<br>(AAC443581; 2.9e-24)      | resistance<br>mechanism                     |

|                          |                 |                 | -10-     |   |                             |
|--------------------------|-----------------|-----------------|----------|---|-----------------------------|
| evbC<br>length<br>319aa  | (55,13556,094)* | (56,10056,103)* | 88, 89   | membrane pump, homolog<br>mithramicin resistance<br>(AAC44357; 1.0e-69)   | resistance<br>mechanism     |
| evbC2<br>length<br>198aa | (56,18456,813)* |                 | 90, 91   | ankrylin like<br>(AAC44356; 0.0041)   | resistance                  |
| evbD<br>length<br>582aa  | (56,96158,709)  | (56,94756,951)  | 92, 93   | acyl-CoA carboxylase<br>(CAB07068; 7.3e-201)  | malonyl-CoA<br>biosynthesis |
| evbE<br>length<br>479aa  | (58,87360,312)  |                 | 94, 95   | IMP dehydrogenase<br>(CAA15452; 4.1e-165)   | tailoring                   |
| evbF<br>length<br>185aa  | (60,47261,029)* | (61,03861,040)* | 96, 97   | hypothetical protein<br>Rv0653c, mycobacterium<br>(CAB07128; 3.8e-06)   | regulator                   |
| evbF1<br>length<br>90aa  | (61,28861,560)  |                 | 98, 99   | unknown   | unknown                     |
| evbF2<br>length<br>152aa | (61,61062,069)  | (61,59761,599)  | 100, 101 | ORFI Streptomyces peucetius (CAA06602; 0.024)   | regulatory/<br>resistance   |
| evbG<br>length<br>557aa  | (62,12263,795)  |                 | 102, 103 | ABC transporter<br>(Q11046; 2.7e-170)   | drug resistance             |
| evbH<br>length<br>645aa  | (63,89165,828)  | (63,88463,887)  | 104, 105 | ABC transporter<br>(Q11047; 5.6e-166)   | drug resistance             |
| evbI<br>length<br>467aa  | (66,46967,872)* | (67,88367,886)* | 106, 107 | lipoamide dehydrogenase<br>(CAA17075; 1.6e-140)   | tailoring                   |
| evbJ<br>length<br>151aa  | (67,97968,434)  |                 | 108, 109 | hypothetical protein<br>Rv3304 [Mycobacterium<br>tuberculosis]<br>(CAA17076; 7.6e-40)                                 | unknown                     |
| evbK<br>length<br>321aa  | (68,52969,494)  |                 | 110, 111 | protease synthase and<br>sporulation regulator;<br>homology to resistance<br>proteins streptomyces<br>(029729; 7.3-7) | regulatory                  |
| evbL<br>length<br>249aa  | (69,61070,359)* |                 | 112, 113 | acetyltransferase/<br>phosphotransferase  | tailoring                   |
| evbM<br>length<br>306aa  | (70,36571,285)* |                 | 114, 115 | hypothetical protein Rv<br>1584c [Mycobacterium<br>tuberculosis]<br>(CAB09085; 0.32)                                  | unknown                     |
| evbN<br>length<br>209aa  | (71,28971,918)* | (71,92671,929)* | 116, 117 | hypothetical protein<br>SC3A7.08 [S. coelicolor]<br>(CAA20071; 4.0e-40)   | unknown                     |
| evbO<br>length<br>230aa  | (72,28472,979)  |                 | 118, 119 | putative lipoprotein [S. coelicolor] (CAA19252; 2.6e-20)  | unknown                     |
| evbP<br>length<br>420aa  | (72,93374,195)* |                 | 120, 121 | peptidase<br>(CAA17077; 6.5e-88)  | unknown                     |

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|                                  |                 |                | -11-     |  |                                |
|----------------------------------|-----------------|----------------|----------|--|--------------------------------|
| evbQ<br>length                   | (74,70776,290)* |                | 122, 123 | methylmalonyl-Coa mutate (BAA30410; 1.8e-149)  | acyl precursor<br>biosynthesis |
| 527aa<br>evbR<br>length<br>696aa | (76,62278,712)  |                | 124, 125 | protein serine/threonine<br>kinase note eukaryotic type<br>(BAA32455;<br>1.1e-71)                                  | regulatory                     |
| evbS<br>length<br>576aa          | (78,79180,521)  |                | 126, 127 | phosphomannomutase<br>(CAA17080; 5.4e-91)  | sugar<br>biosynthesis          |
| evbT<br>length<br>286aa          | (82,07382,933)  |                | 128, 129 | hypothetical protein<br>SC5C7.22c<br>(CAA20634; 5.7e-28)   | 10-28                          |
| evbU<br>length<br>202aa          | (83,28083,888)* |                | 130, 131 | glucose-6-phosphate 1-<br>dehydrogenase low<br>BLAST homology<br>(S61167; 0.00039)                                 | unknown                        |
| evbV<br>length<br>193aa          | (84,08084,661)* |                | 132, 133 | uracil phosphoribosyl<br>transferase<br>(CAA17081; 5.6e-60)  | unknown                        |
| evbW<br>length<br>338aa          | (84,89085,906)* |                | 134, 135 | deoxyribose-phosphate<br>aldolase<br>(AAA79343; 1.3e-54)   | unknown                        |
| evbX<br>length<br>477aa          | (85,90987,342)  |                | 136, 137 | aldehyde dehydrogenase<br>(AAB84440; 4.2e-103)   | tailoring                      |
| evbY<br>length<br>245aa          | (87,42288,159)  | (87,40787,411) | 138, 139 | aldehyde dehydrogenase<br>(CAA71003; 3.4e-16)  | tailoring                      |
| evbZ<br>length<br>137aa          | (88,29288,705)  | (88,28088,282) | 140, 141 | hypothetical protein<br>(CAB06141; 1.3e-16)  | unknown                        |
| evcA<br>length<br>301aa          | (88,71689,621)  |                | 142, 143 | hypothetical protein,<br>putative integral membrane<br>protein [Streptomyces<br>coelicolor]<br>(CAB06143; 4.5e-28) | unknown                        |
| evcB<br>length<br>416aa          | (89,81791,067)  |                | 144, 145 | cytochrome D oxidase<br>subunit I<br>(P94364; 3.0e-65)   | tailoring                      |
| evcC<br>length<br>335aa          | (91,07892,085)  | (91,06891,072) | 146, 147 | cytochrome D oxidase<br>subunit II<br>(CAA71118; 1.9e-15)  | tailoring                      |
| evcD<br>length<br>561aa          | (92,14893,833)  |                | 148, 149 | ABC transporter<br>(CAA22219; 2.6e-107)  | resistance                     |
| evcE<br>length<br>613aa          | (93,83095,671)  |                | 150, 151 | ABC transporter<br>(AAC44070; 3.4e-32)   | resistance                     |
| evcF<br>length<br>229aa          | (95,72996,418)  |                | 152, 153 | unknown  | unknown                        |
| evcG<br>length<br>111aa          | (96,44096,775)* |                | 154, 155 | unknown<br>(AAB84787; 1.9e-8)  | unknown                        |
| evcH<br>length<br>303aa          | (96,89497,805)  |                | 156, 157 | unknown<br>(CAA17083; 9.2e-5)  | unknown                        |

|                                    |                  |                      | -12-     |   |            |
|------------------------------------|------------------|----------------------|----------|---|------------|
| evcI<br>search<br>length<br>691aa  | (98,287100,362)  |                      | 158, 159 | unknown<br>(CAA19992; 6.0e-6)   | unknown    |
| evcJ<br>length<br>197aa            | (100,733101,326) |                      | 160, 161 | putative ATP/GTP binding<br>protein<br>(CAA19989; 7.9e-59)                                    | unknown    |
| evcJ2<br>length<br>134aa           | (101,328101,732) |                      | 162, 163 | unknown<br>(CAA19986; 8.6e-23)  | unknown    |
| evcK<br>length<br>117aa            | (101,803102,156) |                      | 164, 165 | unknown<br>(CAA19991; 1.7e-36)  | unknown    |
| evcL<br>search<br>length<br>1145aa | (102,204105,641) |                      | 166, 167 | unknown<br>(CAA19992; 4.6e-99)  | unknown    |
| evcM<br>length<br>201aa            | (105,907105,641) |                      | 168, 169 | putitive uridine kinase<br>(CAA19591; 1.0e-9)   | unknown    |
| evcN<br>length<br>358aa            | (106,513107,589) |                      | 170, 171 | unknown<br>(CAA17085; 7.5e-120)   | unknown    |
| evrMR<br>length<br>320aa           | (107,653108,615) | (107,637107,641<br>) | 172, 173 | homology to 23S rRNA<br>methylase for mycinamicin<br>resistance (myrA)<br>(BAA03674; 1.4e-79) | resistance |
| evrMR2<br>length<br>193aa          | (108,635109,216) |                      | 174, 175 | homology to gene linked to myrA   | resistance |

# <u>Table 1b. Gene Products and Putative Enzymatic Functions</u> Involved in Everninomicin Production

| ORF1 length | (189-1064)*  | (1069-1073) | 183, 184 | Transcriptional regulator   | unknown      |
|-------------|--------------|-------------|----------|-----------------------------|--------------|
| 291aa       |              |             |          | Biotinylation H70979; 8e-31 | 1            |
| ORF2 length | (1184-2767)* |             | 185, 186 | Propionyl-CoA carboxylase   | unknown      |
| 527aa       |              | 1           |          | T42208; 0.0e                |              |
| ORF3 length | (2863-3753)* |             | 187, 188 | unknown                     | unknown      |
| 296aa       |              |             |          |                             |              |
| ORF4 length | (3776-4276)* | (4280-4284) | 189, 190 | ECF sigma factor T36644;    | regulation   |
| 166aa       |              | `           |          | 8e-26                       |              |
| ORF5 length | (4526-5368)* |             | 191, 192 | Membrane protein            | unknown      |
| 280aa       |              |             |          | CAB94598.1; 5e-50           |              |
| ORF6 length | (5392-6147)* | (6152-6156) | 193, 194 | rRNA methyltransferase      | resistance   |
| 251aa       |              | , ,         |          | AAG32067.1; 4e-49           |              |
| ORF7 length | (6194-7282)* |             | 195, 196 | O-methyl transferase        | modification |
| 362aa       |              |             | ,        | PP42712; 4e-59              |              |
| ORF8 length | (7280-8133)  | (8141-8145) | 197, 198 | unknown                     | unknown      |
| 284aa       |              | `           | •        |                             |              |

| ORF9 length<br>354aa  | (8254-9318)         | (9324-9328) | 199, 200 | oxidoreductase<br>AAG05128.1; 3e-51        | modification          |
|-----------------------|---------------------|-------------|----------|--|-----------------------|
| ORF10 length<br>309aa | (9575-10,504)       | (9568-9571) | 201, 202 | unknown                                    | unknown               |
| ORF11<br>Length 333aa | (10,584-<br>11,585) |             | 203, 204 | deoxyhexose ketoreductase<br>T17473; 1e-49 | sugar<br>modification |

#### Legend for Tables 1a and 1b

- \* CDS, RBS complement on full length biosynthetic locus sequence
- <sup>1</sup> CDS is then putative coding sequence.
- <sup>2</sup> RBS is the putative ribosome binding site.
- <sup>3</sup> GenBank protein database (http://www.ncbi.nih.gov/Entrez/protein.html)
- <sup>4</sup> The first number corresponds to the nucleotide sequence and the second number corresponds to the amino acid sequence.

Although the term "enzymes" is used to refer to the EV biosynthetic pathway gene products, such gene products may be proteins with non-enzymatic functions. Such proteins are also contemplated as falling within the scope of the present invention.

An "EV biosynthetic pathway bottleneck gene" is a gene encoding a product whose level limits the rate of synthesis of everninomicin. Examples of such gene products include, though are not limited to, evrJ (involved in orsellinic acid biosynthesis); evrV, evrW, and evrX (involved in dTDP-glucose synthesis); evbD (involved in malonyl-CoA-synthesis, which is required for orsellinic acid synthesis); and oxidases responsible for oxidation of the amino group on the terminal sugar to produce everninomicin that contains a nitrososugar group. Other likely bottleneck genes include those encoding glycosyltransferases (evdD, evdF, evdH, evdL, and evrS) and tailoring enzymes, particularly sugar modification enzymes.

A modified *Micromonospora carbonacea* refers to a microorganisms that has been genetically engineered to over-express or suppress expression of an EV biosynthetic pathway gene product (enzyme). Such genetic engineering and manipulation is described in detail, *infra*. Preferably, to increase the level of production of everninomicin, the modified microorganism overexpresses one or more bottleneck genes. To produce an everninomicin analog or homolog, various tailoring enzyme genes (*e.g.*, evdB, a hexose aminotransferase that produces an amino sugar; evrF, a nonheme halogenase that chlorinates the orsinillic acid; or an oxidase gene that produces a nitrososugar by oxidation of an aminosugar) may be knocked out. Other knock-outs may be made of putative key genes, resulting in all likelihood in blockage of everninomicin biosynthesis. These include the orsellinic acid synthase (evrJ), dTDP-glucose synthases (evrV, evrW, and evrX), and glycosyltransferases (evdD, evdF, evdH, evdL, and evrS). A knockout of the glycosyltransferase that adds the terminal glycosyl group is expected to produce an everninomicin analog lacking the terminal glycosyl group.-



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### Please replace the star agraph of page 4 (lines 20-22) with the following paragraph:

- In addition, isolated polypeptides corresponding to an everninomic biosynthetic pathway gene product are provided. Specific open reading frames and amino acid sequences of the polypeptides are set forth in Figure 11 (SEQ ID NOS: 2-175) and Figure 12 (SEQ ID NOS: 183-204).

## Please replace the 7<sup>th</sup> full paragraph of page 5 (lines 24-25) with the following paragraph:

-- Figure 7A-B. (A) Map of pSPRH840 integrating vector. (B) Sequence of integrase gene (SEQ ID NO: 176) and deduce amino acid (SEQ ID NO: 177).--

## Please replace the 11<sup>th</sup> full paragraph of page 5 (lines 32-33) with the following paragraph:

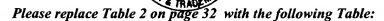
--Figure 11A-AB. Everninomic biosynthetic pathway locus sequence (SEQ ID NO:1) with open reading frames and deduced amino acid sequences (SEQ ID NOS: 2-175).--

## Please replace the 12th full paragraph of page 5 (lines 34-35) with the following paragraph:

--Figure 12A-G. Everninomic biosynthetic pathway locus sequence (SEQ ID NO: 182) with open reading frames and deduced amino acid sequences (SEQ ID NOS: 183-204).--

#### Please replace the 3<sup>rd</sup> full paragraph of page 14 (lines 16-20) with the following paragraph:

--A *Micromonospora* site-specific Att/Int functions consist of an integrase protein and AttP site, *e.g.*, as depicted in Figure 7B (SEQ ID NO: 177) and in a specific embodiment encoded by a nucleic acid having a sequence as depicted in Figure 7B (SEQ ID NO: 176), that permits site-specific integration of a vector into an actinomyce, and particularly a *Micromonospera*, genome.--



-- Table 2. Orsellinic Acid Biosynthetic Gene Products

| Gane<br>Product | CDS           | aroidizo | Birgymedelfinedań  |
|-----------------|---------------|----------|--|
| evrF            | 21,06422,542  | 36,37    | non-heme   |
|                 |               |          | oxygenase/halogenase addition                            |
| evrI            | 25,55026,626  | 42,43    | acyl starter unit  |
| evrJ            | 26,68530,479  | 44,45    | Orsellinic acid synthase/6-methylsalicilic acid synthase |
| evbD            | 56,96158,709  | 92,93    | acyl-CoA carboxylase                                     |
| evbQ            | 74,70776,290* | 122,123  | Methylmalonyl-CoA mutase                                 |

#### Please replace Table 3 on page 34 with the following Table:

-- Table 3. Sugar Biosynthetic Gene Products

| Table 3. Sugar Blosynthetic Gene Products |                                  |         |                                 |  |
|---|----------------------------------|---------|---------------------------------|--|
| Com.<br>Raduet                            | If the the state is the first of | SEQUING | Dizzinetle Dinefon              |  |
| evdA                                      | 1321382*                         | 2, 3    | Hydroxylase                     |  |
| evdB                                      | 14902611*                        | 4, 5    | hexose aminotransferase         |  |
| evdC                                      | 26223860*                        | 6, 7    | oxidase (flavoprotein)          |  |
| evdE                                      | 53096235                         | 10, 11  | hexose dehydratase              |  |
| evdI                                      | 946310,224*                      | 18, 19  | Hydrolase                       |  |
| evdK                                      | 11,20812,455                     | 22, 23  | hexose dehydratase or epimerase |  |
| evrA                                      | 14,41015,363*                    | 26, 27  | hexose epimerase                |  |
| evrB                                      | 15,38016,414*                    | 28, 29  | hexose oxidoreductase           |  |
| evrC                                      | 16,41917,873*                    | 30, 31  | hexose dehydratase              |  |
| evrD                                      | 17,87018,934*                    | 32, 33  | GDP-mannose 4,6-dehydratase     |  |
| evrV                                      | 41,67942,707*                    | 68, 69  | dTDP-glucose epimerase          |  |
| evrW                                      | 42,81043,799*                    | 70, 71  | dTDP-glucose dehydratase        |  |
| evrX                                      | 43,79944,866                     | 72, 73  | dTDP-glucose synthetase         |  |

#### Please replace the remainder of Table 3 on page 35 with the following Table: --

| evbS  | 78,79180,521 | 126, 127 | Phosphomannomutase        |
|-------|--------------|----------|---------------------------|
| evbU  | 83,28083,888 | 130, 131 | Glucose-6-phosphate 1-    |
|       |              |          | dehydrogenase             |
| ORF9  | 82549318     | 199, 200 | Oxidoreductase            |
| ORF11 | 10,58411,585 | 203, 204 | Deoxyhexose ketoreductase |



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Please replace Table 4 on page 35 with the following Table:

-- Table 4. Glycosyltransferases

| Gene :<br>Product | CDS           | SEQ IID No. | Rmzymatic Function              |
|-------------------|---------------|-------------|---------------------------------|
| evdD              | 41435312      | 8, 9        | DNTP-hexose glycosyltransferase |
| evdF              | 62327275      | 12, 13      | DNTP-hexose glycosyltransferase |
| evdH              | 83429364      | 16, 17      | DNTP-hexose glycosyltransferase |
| evdL              | 12,10813,022* | 24, 25      | DNTP-hexose glycosyltransferase |
| evrS              | 38,89240,163* | 62, 63      | DNTP-hexose glycosyltransferase |

Please replace Table 5 on page 36 with the following Table:

-- Table 5. Tailoring Gene Products

| - Table 5. Tailoring Gene Products |               |            |                                      |  |  |
|------------------------------------|---------------|------------|--------------------------------------|--|--|
| Gene Product                       | CDS           | SEQ ID No. | Enzymatic Function                   |  |  |
| evrG                               | 22,74824,172  | 38, 39     | oxidase                              |  |  |
| evrL                               | 31,94132,882* | 48, 49     | heme biosynthesis                    |  |  |
| evrM                               | 33,16734,405* | 50, 51     | p450 hydroxylase                     |  |  |
| evrN                               | 34,44935,210* | 52, 53     | methyl transferase                   |  |  |
| evrQ                               | 36,99838,026* | 58, 59     | oxidoreductase/heat stress protein   |  |  |
| evrT                               | 40,21640,890  | 64, 65     | L-proline hydroxylase                |  |  |
| evrU                               | 40,88741,576  | 66, 67     | methyltransferase                    |  |  |
| evbA                               | 53,55454,207  | 84, 85     | o-methyltransferase                  |  |  |
| evbE                               | 58,87360,312  | 94, 95     | IMP dehydrogenase                    |  |  |
| evbI                               | 66,46967,872* | 106, 107   | lipoamide dehydrogenase              |  |  |
| evbL                               | 69,61070,359* | 112, 113   | acetyltransferase/phosphotransferase |  |  |
| evbX                               | 85,90987,342  | 136, 137   | aldehyde dehydrogenase               |  |  |
| evbY                               | 87,42288159   | 138, 139   | aldehyde dehydrogenase               |  |  |
| evcB                               | 89,81791,067  | 144, 145   | cytochrome D oxidase subunit I       |  |  |
| evcC                               | 91,07892,085  | 146, 147   | cytochrome D oxidase subunit II      |  |  |
|                                    |               |            | 1                                    |  |  |



Please insert Table Property page 37 with the following Table: --

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Table 6. Regulatory Gene Products

| Core //<br>Product | CDS             | SEO ID KO | Dissinate Dungton  |
|--------------------|-----------------|-----------|--|
| evrR               | 38,07238,566    | 60, 61    | hexaheme nitrite reductase regulator/<br>methyltransferase |
| evsA               | 47,15649,234*   | 78, 79    | serine-threonine kinase                                    |
| evbF               | 60,47261,029*   | 96,97     |  |
| evbF2              | 61,61062,069    | 100, 101  |  |
| evbK               | 68,52969,494*   | 110, 111  | protease synthase/sporulation regulator                    |
| evbR               | 76,62278,712    | 124, 125  | protein serine-threonine kinase<br>(eukaryotic type)       |
| evcJ               | 100,733101,326* | 160, 161  | ATP/GTP binding protein                                    |
| ORF1               | 1891064*        | 183, 184  | Transcriptional regulator biotinylation                    |
| ORF4               | 37764276*       | 189, 190  | ECF sigma factor   |

## Please replace Table 7 on page 37 with the following Table:

- - Table 7. Resistance Mechanism Genes

| Table 7. Resistance Weenanism Genes |               |          |                             |  |  |
|-------------------------------------|---------------|----------|-----------------------------|--|--|
| Gene Proc                           | lier CDS      | SEQUENCE | Janayne ile littre ion      |  |  |
| evrE                                | 19,37420,906  | 34, 35   | multidrug eflux transporter |  |  |
| evrY                                | 45,01445,760* | 74, 75   | dehalogenase                |  |  |
| evrZ                                | 45,96246,714* | 76, 77   | muramidase/lysozyme         |  |  |
| evbB                                | 54,36255,117* | 86, 87   | membrane pump               |  |  |



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#### Please replace the remainder of Table 7 on page 38 with the following Table: --

| evbC   | 55,13556,094*  | 88, 89   | membrane pump          |
|--------|----------------|----------|------------------------|
| evbC2  | 56,18456,813*  | 90, 91   | ankrylin-like          |
| evbG   | 62,12263,795   | 102, 103 | ABC transporter        |
| evbH   | 63,89165,828   | 104, 105 | ABC transporter        |
| evcD   | 92,14893,833   | 148, 149 | ABC transporter        |
| evcE   | 93,83095,671   | 150, 151 | ABC transporter        |
| evrMR  | 107,653108,615 | 172, 173 | 23S rRNA methylase     |
| evrMR2 | 108,635109,216 | 174, 175 |                        |
| ORF6   | 53926147*      | 193, 194 | rRNA methyltransferase |

#### Please replace the 1<sup>st</sup> full paragraph of page 44 (lines 6-13) with the following paragraph:

- In a specific embodiment, the present invention relates to a new nucleic acid sequence, to vectors for its expression and to its use in fermentation processes in actinomycetes. This nucleic acid sequence encodes a *Micromonospera*, and particularly *M. carbonacea*, *var. africana*, att/int functions and thus permits development of an integrating vector. In a specific embodiment, the att/int functions has an amino acid sequence as depicted in SEQ ID NO: 177. In a more specific embodiment, the integrase is encoded by a nucleic acid having a nucleotide sequence as depicted in SEQ ID NO: 176 (Figure 7B). A preferred integrating plasmid is shown in Figure 7A.- -

#### Please replace the last full paragraph on page 48 with the following:

- - The coding region of *evrF* gene was amplified with PCR primers:

5' PR 657 CCC TCG AGA TGT CCA GCA AGA TCC TA (SEQ ID NO: 178);

3' PR 658 CGA ATT CTC AGG CAG ACT GCT CTG (SEQ ID NO: 179); and

5' PR 659: CCC TCG AGA ATG TCC AGC AAG ATC CTA (SEQ ID NO: 180);

3' PR 660: CGA ATT CAG ACT GCT CTG CCG CCG C (SEQ ID NO: 181);

using the Advantage-GC Genomic PCR kit and Advantage HF polymerase (Clontech, Palo Alto, CA) and a Perkin-Elmer 9600 PCR machine (Foster City, CA). The 1.5 kb PCR products were digested with *XhoI* and *EcoRI* and the fragments were ligated to *XhoI* and *EcoRI* digested pBADHisA (primer pair PR657/PR658 product) and pBADMycHisC (primer pair PR659/PR660 product) and transformed into *E. coli* Top10 (Stratagene, LaJolla, CA). Transformants were analyzed by plasmid isolation followed by digestion and gel electrophoresis analysis. Appropriate clones were also verified by sequence analysis. This yielded the evrF expression clones pSPRE59 (pBADHisA) and pSPRE19 (pBADMycHisC). Top10 cells containing either pSPRE59 and pSPRE19 were grown overnight at

 $37^{\circ}$ C with shaking in LB containing 50ug/ml AMP. Overnight cultures were used to innoculate fresh LB containing 50 µg/ml and grown at 37 °C with shaking to an OD<sub>600</sub> of 0.4 to 0.5. L-arabinose was added to a final concentration of 0.02% and the culture was incubated for an additional 4 hours. Cells were collected by centrifugation, resuspended in 100 µl Tris-Glycine buffer and boiled for five minutes. Whole cell protein lysate was loaded onto a SDS-PAGE gel, electrophoresed, and stained with coomassie blue to determine protein expression.- -